

### 0000-2952(95)00160-5

# **Short Communication**

ABSORPTION, TISSUE DISTRIBUTION AND IN VEO STABILITY IN RATS OF A HYBRID ANTISENSE OLIGONUCLEOTIDE FOLLOWING ORAL ADMINISTRATION

SUDHIR AGRAWAL,\*\* XUESHU ZHANG.; ZHIHONG LU., HUI ZHAO.;
JEFFREY M. TAMBURIN.; HEMING YAN; HONGYING CAL.)
ROBERT B. DIASIO.; IVAN HABUS.\* ZHIWELHANG.\*
RADHAKRISHNAN P. IYER.\* DONG YU\* and RUIWEN ZHANG.;

\*Hybridon, Inc., Worcester, MA 01605; and ‡Department of Pharmacology and Toxicology, and Division of Clinical Pharmacology, University of Alabama at Birmingh in Birmingham, A1, 35294 U.S.A.

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Abstract—In vico stability and oral bioavailability of an oligodoxy, indefeotide pho phorothioare containing segments of 2°-O-methyloligoribonucleotide phosphorothioates at both the 3°- and 5°-ends (hybrid oligonacleotide) were studied. A 25-mer "S-labeled hybrid oligonacleotide was administered to rats by gavage at a dose of 50 mg/kg body weight. HPLC analysis revealed that this hybrid oligonucleotide was stable in the gastrointestinal tract for up to 6 hr following oral administration Radioactivity associated with the hybrid oligonucleotide was detectable in portal venous plasma systemic plasma, various tissues, and urine. Intact hybrid oligonucleotide was detected, by HPLC analysis in portal venous plasma, systemic plasma, and various tissues. The majority of the radioactivity in urine was associated with degradative products with lower molecular weights, but the intact form was also detected. In summary, the hybrid oligonucleotide was absorbed intact through the gastrointestinal tract, indicating the possibility of oral administration of oligonucleotides, a finding that may be important in the development of antisense oligonucleotides as therapeutic agents.

Key nords, antisense obgonucleotides; HIV; oral bioavailability

The use of antisense ofigonacleotides has been shown to be a promising approach in the development of therapeutic agents for the treatment of viral infections (including HIVs intection) and cancer [1]. In fact, several unmodified and modified antisense oblgodeoxynucleotides have been demonstrated to have antisense activity, both in vitro and in vivo [1]. Thus fat, most in vivo biological and pharmacokimetic studies have been carried out with PS-ofigonucleotides [2–10]. In general, PS-ofigonucleotides have a short distribution half-lift and a longer elimination half-lift in plasma, and are distributed widely into all major tissues following a vinjection [2–8, 10]. Varying in vino degradation of PS-ofigonucleotides has been observed [2–10]. Several antisense ofigonucleotides recently have entered early climal evaluations including hum in

pharmacokinetic studies [9,1]. Compared with the results from annual studies. PS oligonucleotides have a similar pharmacokinetic profile in humans (9, 1).

Attempts have been made recently to stabilize PS-obgonucleotides to avoid degradation in vivo. It has been observed that 2°-O-methylologoribonucleotide phosphorothioates are more resistant to nucleases than the PS-obgonucleotide in viiro [11-13]. By incorporating segments of 2°-O-methylologoribonucleotide phosphorothioate at the ends of PS-obgonucleotide, a significant increase in protection against exonuclease has been observed in viiro. In viirous assay models obgonucleotide phosphorothioate observed in obgoniomic office phosphorothioates at both the 3°- and 5°- ends (hybrid of souncleotide) I as been shown to have greater stability in citic and in come, as well as improved antiscusse activities as a PS obgonic cotides [41, 12]

More recently at two thosto-only, disposition, and excittion of a 25 mer hybrid obsonucleotide were determined in rate there is, bone administration of 38 labeled hybrid on, oracleotide at a dose of 91 mg kg [14]. The plasma disappearance curve for the hybrid obsonucleotide count to described by a two-compartmental model, with half most of 6.34 and 52.02 hr. The majority of radioactivity in plasma was associated with the infact hybrid obsonucleotide. Urmary corretion represented the major pathway of chanation, and fecal exerction was a minor pathway of chanation. A wide tissue distribution of hybrid obsonucleotide was observed based on radioactivity levels and analysis by HPTC showed that the majority of the rathoactivity was issociated with the intact hybrid object in toolkie. Compared with other oil

Corresponding authors S.A. Tel. (508) 751-7531, FAX (508) 781-7692. R.Z. Tel. (205) 934-8888, FAX (205) 934-8240.

<sup>§</sup> Abbreviations HIV, human immunodeficiency virus, PS ofigonucleotide oligodeoxynucleotide phosphorothoate, and hybrid oligonucleotide hybrid oligonucleotide phosphorothoate.

Zhang R. Yan J. Shahiman H. Anim G. Lu Z. Saag MS, Jiang Z. Temsamani J. Schechter PJ. Martin RR. Agrawal S and Drasio RB. Haman pharmacokinetics of an anti-HIV antisense oligonucleotide phosphorothioate (CLM91) in HIV intected individuals. The 34th Interscience Conference on Antimicrobial Agents and Chemomerapy, Orlando, FL. Oct. 4.7, 1994. Later Breaker Session AS, p. 2.

which have a similar nucleotide sequence [8], the hybrid ofigonucleotide has been found to have a shorter distribution half-life and a longer elimination half-life, and is more stable in various tissues [14]. More interestingly, following to impection of this hybrid ofigonucleotide to rats, infact compound was detected in the liver and intestinal tissue and contents, indicating that there is an enterotiepatic circulation of this objoinucleotide [14]. These results suggest the possibility of alternative delivery of antisense ofigonucleotides as the appearance agents in the future. The purpose of the present study was to determine absorption, tissue distribution, and in vivo stability of the 25-mer nyorid ofigonucleotide in rats following or al administration

# Materials and Methods

Synthesis of ionlabeled and \*S-labeled hybrid oligonucleotide. The 25-mer hybrid oligonucleotide (5-CUCU-CGCACCCATCTCTCTCC-UUCU-3') contains seaments of four 2'-O-methyloligoribonucleotide phosphorothicates at both the 3'- and 5'-ends. Synthesis of the hybrid origonucleotide was carried out using deoxynucleoside phosphoramidites (Milligen, Milford, MA) and 2'-O-methylribonucleoside phosphoramidites (Glen Research, Sterling, VA) on an automated synthesizer (Biosearch 8800). The synthesis was carried out on a 1 mM scale by using the protocols reported earlier [11, 14, \*] The purity of the hybrid oligonucleotide was shown to be greater than 90% by analyses of "P NMR, ion exchange HPLC, polyacrylamide gel electrophoresis and melting temperature (both UV and circular dichroism), with the remainder being n-1 and n-2 products. To prepare the ' labeled hybrid ongonucleotide, synthesis was carried out in a way similar to that described above except that the last five couplings were carried out using 2'-Omethylribonucleoside H-phosphonate. The 2'-O-methylhonucleoside H-phosphonates, U and C, were synthesized by following the PCL/triazole method,\* starting from the appropriate 2%O methylribonucleoside, and isolated as thethylammonium salts. The isolated 2'-O-methylribonucleoside H phosphonates, U and C, were analyzed by both "P and H NMR spectroscopies. Prior to use in the oligonucleotide synthesis, the nucleoside  $H_{\gamma}$ phosphonates were evaporated to dryness twice with anhydrous pyridine and dissolved into anhydrous pyridine CH.CN (1.1) to a concentration of 40 mM. After assembly, the controlled pore glass (CPG)-bound obgonucleotide containing four H-phosphonate linkages was oxidized with "S elemental sulfur (Amersham; 0.5 to 2.5 Ci/militatom) and deprotected by the same procedure as reported earlier [11] Purification of the 38-labeled hybrid oligonucleotide was carried out by PAGE (20%). 7 M urca). The visualized band product under UV light was excised, extracted in 100 mM ammonium acetate, and desatted using a Sep-Pak C18 column (Waters, Millord, MA) The specific activity of the hybrid oligonucleotide obtained was 0.25 aCi/ug. The purity of the 2S-labeled hybrid origonic leotide was shown to be greater than 98%. by analyses of "P NMR- ion exchange HPLC, and PAGE with the remainder being n-1 and n-2 products.

Animals and treatment. The protocol for animal use and care was approved by the Institutional Animal Care and Use Committee of the University of Alabama at Birmingham Male Sprague-Dawley rats (110 ± 10 g; Harlan Laboratories, Indianapolis, IN) were utilized. The animals were given commercial diet and water ad lib. for I week prior to the study. Unlabeled and <sup>38</sup>S-labeled

offigural actions where a described in physical great values power, Sach is a concentration of 25 mg ml and were administrated to the animals via gavage at a dose of 50 mg. kg (sp. act. 12.40 mg). Doses were based on the preticalized tody weight and rounded to the nearest 0.01 ml.. After dosing, each animal was placed in a metabolism cago and fed a commercial diet and water ad ho. Fotal voided urine was collected, and each metabolism cage was washed following the collection intervals. Total excreted teccs was collected from each animal at various time points, and feces samples were homogenized prior to quantitation of radioactivity. Blood samples were collected and the animals were killed at various times (i.e. 1, 3, 6, 12, 24, and 48 far, 2 rats/time point). Plasma was separated by contribugation. Following removal, tissues/organs were trimmed of extraneous fat or connective tissue, emptied and cleaned of all contents, and individually weighed prior to homogenization. To determine the chemical form(s) of radioactivity is the portal venous blood, a separate group of animals was subjected to the same treatment procedure as above. Portal venous blood samples (1/rat) were taken at scheduled times (1, 3, and 6 hr), and plasma was separated by contribugation

To quantitute the total absorption of the hybrid ofigonecleorade two additional groups of animals (3/group) were treated using the same procedure as above. Animals were killed at 6 or 12 hr post-dosing, and the gastrointestinal tract was then removed. Radioactivities in the gastrointestinal tract, feces, urine, plasma, and the remainder of the body were determined separately. Total recovery of radioactivity was also determined to be 95 ± 6%. The percentage of the absorbed hybrid ofigonucleotide-derived radioactivity was determined by the following calculations.

(total radioactivity in the remainder of the body and plasma + total radioactivity in urine)

ttestal radioactivity in the gastrointestinal tract, to see, urine plasma, and the remainder of the body)

Sample preparation and total radioactivity measurements. The total radioos bythes in tissues and body fluids were determined by liquid scintillation spectrometry (LS 0000) FA; Beskman, fryine, CA), using a method described previously [5, 141]. In biret, biological fluids (plasma, \$0–100 aL; urine, \$0–100 aL), were mixed with 6 mL of scintillation solvent (Beckman) to determine total radioactivity. Feese were ground and weighed prior to being homogenized in a 9-told volume of 0.995 NaCl. Following their removal, tissues were blotted immediately on Whatman No. 1 filter paper and weighed prior to being homogenized in 0.995 NaCl (3.5 mL) givet weight). An aliquot of the homogeniate (100 aL) was mixed with solubilizer (18-2; RPI, Mt. Prospect, IL) and then with scintillation solvent (6 mL) to permit quantitation of total radioactivity.

HFT Canalysis. The radioactivities in plasma and urine samples were analyzed by ion-paired HPLC using a modification of a method described previously [4]. Unine samples were centrifuged and passed through a 0.2 µm Acre blief (Gelman, Ann Arbor, MI) prior to analysis Hyland obgenucicotide and metabolites in plasma and tissue samples were extracted by methods described previously [14] Using the extraction methods, the extraction effusioney was approximately 75% for 18-25 mer and our. Tor \$ 10 mer. Monomer cannot be extracted using this method. A Microsorb MV-C4 column (Rainin Instruments, Wobarn, MA) was employed in HPLC using a Hewlett-Packard 1050 HPLC with a quaternary pump for gradient making. The mobile phase included two butfers: Butter A was 5 mM PIC-A reagent (Waters Co., Bedford MAs in water and Buffer B was 4:1 (v.v.) acctomatile (1 shor) water. The column was cluded at a flow rate of 1.5 mL, min, using the following gradient! (1) Q Smin, 9% buttet B: (2) \$ 45 min 6-35% Bufter B: and 13, 18 Winn. S 80% Butter B. The column was

<sup>•</sup> Metelev V and Agrawal S, Hybrid oligonucleotide phosphoroilhoates. Synthesis, properties and anti-HIVactivity. Proceedings of International Conferences on Nucleic Acid Medical Applications. Cancun, January 1993. Abstract 1-1

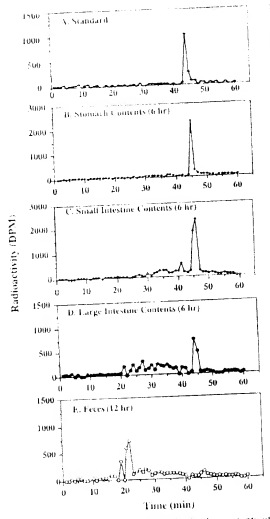


Fig. 1. HPLC analysis of radioactivity in the contents of stomach, small intestine, large intestine, and feees. The radioactivities in plasma samples were analyzed by ion-parted HPLC, using a Microsorb MV-C4 column and gradient elution (see Materials and Methods). Under the conditions used in the present study, the retention time for the standard "S-labeled 25-mer hybrid oligonucleotide was 45.0 min (panel A). This figure shows representative chromatograms for samples from animals killed at 6 hr after dosing. Similar HPLC profiles were observed with each of the animals in the present study.

equilibrated with Buffer A for at least 30 min prior to the next run. By using a RediFrac fraction collector (Pharmacia I KB. Biotechnology. Piscataway. NJ), 1-min. fractions (1.5 mL) were collected into 7-mL scintillation vials and mixed with 5 mL of scintillation solvent to determine the radioactivity in each fraction.

#### Results

In vivo stability of hybrid oligonucleotide in the gastroniesunal tract. The hybrid oligonucleotide was shown to be stable in the stomach and in the small and large intestines (Fig. 1), as analyzed by HPLC. Up to 6 hr totlowing administration, the hybrid oligonucleotide

remained artasy in the stomach and its contents and the majority of cadioacters in the small artestine were associated with the intact ofigonucleotide. Extensive degradation of this ofigonucleotide was observed in the large intestine 6 hi after dosing (Fig. 1)

Absorption of technic oligonic conde following oral administration. Radioactivity was detectable in various tissues following oral administration of the radiolabeled hybrid oligonic leotide. Figure 2A illustrates the plasma, liver, kidney and spleen concentration-time course of the oligonic leotide equivalents after oral administration of radiolabeled organical conde. The hybrid oligonic leotide derived radioactivity was detected in all tissues examined. Figure 2B illustrates the concentration-time courses for several representative tissues. An accumulation of radioactivity was observed in these tissues.

Urmary excretion of hybrid oligonacleotide following oral administration. Radioactivity was detected within 1 hr tollowing oral administration of the hybrid oligonacleotide. Following a complete urine collection, the mean cumulative excretion of urnary radioactivity was determined to be 25% of the administered dose over 24 hr and 3.8% over 48 hr post-dosing. The majority of radioactivity in the urine was associated with the degradative products of the hybrid oligonacleotide. However, trace intact hybrid oligonacleotide was also detected (Fig. 3).

Chemical forms of radioactiony in portal venous blood, systemic plasma, and various tissues following oral administration. The chemical forms of radioactivity in portal venous plasma, systemic plasma, liver, and kidneys were examined turther by HPLC, demonstrating the presence of both intact and degraded products of the hybrid oligonucleotide in these samples (Fig. 4)

Bioaualability of hybrid oligonucleotide following oral administration. No significant degradative products were detected in the stomach contents. The majority of the radioactivity in the small intestine contents was present as the intact compound for up to 12 hr post-dosing. The radioactivity in the large intestine was present as both the intact form and degradative products. Based on the quantitation of total radioactivity in the gastrointestinal tract, feecs, urine, plasma, and the remainder of the body in two groups of animals (3/group), the total absorption of the hybrid oligonucleotide-derived radioactivity was determined to be  $10.2\pm2.5\%$  over 6 hr and  $25.9\pm4.7\%$  over 12 hr following oral administration of the radioactivity in the study was 95.5%.

## Discussion

The rationale of the present study is to demonstrate oral bioavailability of antisense oligonicleotides, which may facilitate the development of this class of compounds as therapeutic agents. The hybrid oligonucleotide was chosen as a test compound because it has shown greater in vitro and in vito stability over other PS-oligonucleotides [11-14. 1 The present study established three major points regarding the oral availability of obgonucleotides: (1) the hybrid oligomicleotide is stable in stomach and small intestine ussues and contents, (2) the hybrid oligonucleotide is absorbed through the portal venous blood system, and (3) the absorbed our nucleotide-derived radioactivity is associated with the intact obgonucleotide and distributed into various tissues, and excreted into urine, as seen previously with the administration [14]. The above data demonstrate oral absorption of oligonic leotide providing

<sup>\*</sup>Metrics V and Agrawal S, Hybrid oligonucleotide phosphorothicates: Synthesis, properties and anti-HIV-activity. Proceedings of International Conferences on Nucleic Acid Medical Applications, Cancun, January 1993, Abstract 1-1

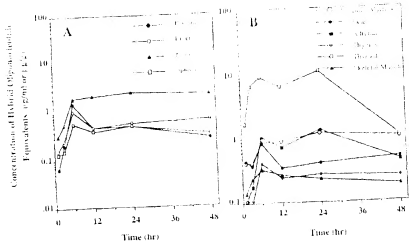


Fig. 2. Plasma and tissue concentration-time courses of radioactivity. Mean plasma and tissue concentrations were expressed as micrograms of hybridized obsolucteotide equivalents per milliliter of plasma of per gram of tissue/organ. Two animals were used for each time point. Dissue concentration was based on the quantitation of radioactivity.

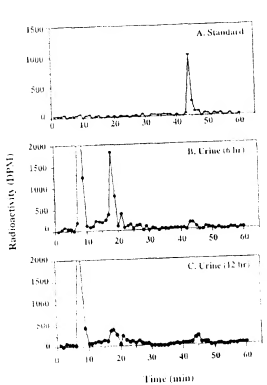


Fig. 3. HPLC analysis of radioactivity in urine. The radioactivities in urine samples were analyzed by HPLC. This figure shows representative chromatograms for urine samples from one animal at various times. Similar HPLC profiles were observed with each of the animals in the present study. Both intact obgonucleotide and metabolites with lower molecular weights were detected in urine with 24 hr post-dosing.

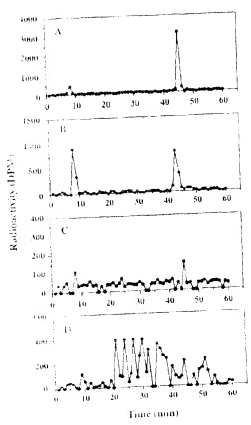


Fig. 1 (iPLC) analysis of radioactivity in the portal venous plasma (A), systemic plasma (B), liver (C), and kidney (D). This figure shows representative chromatograms for biological samples from one animal at 6 hr post-dosing Similar HPLC phofiles were observed with other animals in the present study.

the basis for future studies of our onucleotides as ofally available therapeutic agents

Support for the oral availability of obgonucleondes derives from several observations. First, the stability of the onyonucleotide in the gastrointestinal tract was demonstrated by HPLC analysis Polyacrylamide gel electrophoresis of extracts from these tissues confirmed the results (data not shown). Second, the majority of the radioactivity in the portal verous plasma was associated with the intact ofigonucleotide, although degradative products were also detected. Third, radioactivity in the liver was associated with intact oligonucleotide as well as metabolites, as seen with r.v. injection. Fourth, intact oligonucleotide was found in systemic plasma, indicating that intact oligonucleotide is available to extrahepatic tissues. Fifth, a wide tissue distribution of radioactivity was observed with the intact oligonucleotide detected. Finally, a significant amount of radioactivity was excreted through urine, with intact obgonucleotide detected in the urine

Despite extensive studies on the chemistry, molecular biology, and in ouro biological activity, limited studies on the pharmacology of antisense obgonucleotides have been conducted. To date, most in vivo animal studies have been carried out with PS-oligonucleotides [2-10]. In a preliminary study [2] we reported the pharmacokinetics of a \*5-labeled 20-mer PS-ofigonucleotide targeted to the HIV tat gene in mice via i.p. and i.v. injection. Radioactivity could be detected in most tissues up to 48 hr post-dosing. Similar results were reported by Iversen and colleagues [7, 8] following a study of \*S labeled 27-mer PS-oligonucleotide complementary to the HIV rev gene in mice at an i.p. dose of 50 mg/kg. Cossum et al [3] reported a study of a Chabeled 20 mer PS-oligonucleotide in female rats at an (v. dose of 3.6 mg/kg. High radioactivities were found in the liver, kidney, bone marrow, skin and skeletal muscle In plasma, the radioactivity was extensively bound to proteins. This PS-oligonucleotide was rapidly absorbed after intradermal injection [6] and tissue distribution of the PS-oligonucleotide was comparable to cv. injection. A preliminary study to rabbits and monkeys by Iversen [7] suggested potential species differences in the pharmacokinetics of PS-oligonacteotides. Several PS-oligonucleotides and modified analogs have been studied recently in our laboratories for comprehensive pharmacologic analyses [5, 10, 14].

Pharmacologic study in humans is extremely limited. although several clinical studies of antisense oligonucleotides have been initiated. Recently, Crooke et al. [9] injected a 20 mer "C-labeled PS-oligonucleotide into patients' genital warts at a dose of 1 mg, wart. Radioactivity was present in wart tissues for 72 hr. Systemic absorption was also observed, and 30% of the radioactivity was climinated through expiratory air. Human pharmacokinetics of a "S-labeled 25-mer PS-ofigonucleotide was performed recently in our laboratory." "S-Lubeled GEM 91 was administered to 6 individuals by 2-hi 15 infusion at a dose of 0.1 mg/kg body weight. Plasma disappearance cutves for GEM 91 derived radioactivity could be described by the sum of two exponentials, with half lives (mean 5 EM) of 0.18 (5.0.04) and 26.71 (\* 1 62) hr. The chemical forms of radioactivity in plasma were further evaluated by PAGE and HPLC, demonstrating the presence of both intact GEM 91 and lower molecular

In conclusion, the present study provides initial evidence for the possibility of alternative delivers of antisense oligonucleotides as the apeutic agents. The mechanisms responsible for absorption of the hybrid oligonucleotide have not been defined in the present study. Further studies examining the mechanisms of transport of oligonucleotides in the gastrointestinal tract and liver, first pass effects of the liver, enterohepatic circulation, and formulation of oligonucleotides are currently underway.

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weight nictus less. Unday exploits to represented the major pathway of cimanation of GPM 91, with 49.15 to 80% of the administred door excreted within 24.15 to 80% of the administred door excreted within 24.15 and 70.37 to 75% over 90 to after GLM 91 administration. The radic activity in urine way analyzed by HPLC and shown to be mainly lower molecular weight metabolities indicating an extensive degradation of the PS obsonuelectude.

The mechanisms responsible for the metabolism of objoinacleotides remain unclear. It is believed that the degradations faintisense oligonucleotides may be structure length-, dose-, administration route-, species-, and tissue dependent. Compared with other ofigonucleotides, PS obgonucleotides are more stable in two. The degradation of PS-oligonucleotides is mainly by 3'-exonucleases [2, 4, 5, 7, 8, 10, 14] After 1 v. injection of a 20 mer PSoligonucleotide into mice, there are no significant degradative products in urine [2]. In contrast, after i p injection, 75, 90°, of the PS-obgonucleotide in urine is degraded [2]. Interestingly, chain length extension of the administered PS oligonucleotide was observed in the kidney, liver and intestine. Recently, Cossum et al. [3, 6] reported that "C-labeled PS-oligonucleotide is eliminated mainly through expiratory air, and no nativet PS oligonucleotide is found in the urine or teces. However other studies have shown that urmary exerction represents the major pathway of channation following administration of H- or "S-labeled oligonucleotides to animals [2, 4, 5, 7, 8, 10, 14] and humans.\*

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